EAST Search History

Ref #	Hits Search Query		DBs	Default Operator	Plurals	Time Stamp		
L1	242	(548/340.1).CCLS.	USPAT; DERWENT	•		2007/12/31 12:31		
L2	6	(("6069251") or ("5405988") or ("4939130")).PN.	USPAT; DERWENT	OR	OFF	2007/12/31 12:34		
L3	8	(("6875757") or ("7060697") or ("7064217") or ("7241790")).PN.	USPAT; DERWENT	OR	OFF	2007/12/31 12:42		
L4	2	("20040224941").PN.	US-PGPUB; USPAT; DERWENT	OR	OFF	2007/12/31 12:43		
L5	2	("20050032744").PN.	US-PGPUB; USPAT; DERWENT	OR	OFF	2007/12/31 12:45		
L6	2	("20050023386").PN.	US-PGPUB; USPAT; DERWENT	OR	OFF	2007/12/31 12:47		
L10	1	("20050107447").PN.	USPAT; DERWENT	OR	OFF	2007/12/31 12:49		
L11	1	("20060135786").PN.	USPAT; DERWENT	OR	OFF	2007/12/31 12:51		
L12	1	("20060223866").PN.	USPAT; DERWENT	OR	OFF	2007/12/31 12:53		
L13	1	("20070088002").PN.	USPAT; DERWENT	OR	OFF	2007/12/31 12:54		

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                 FSTA enhanced with new thesaurus edition
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         SEP 17
                 CAplus coverage extended to include traditional medicine
                 patents
                 EMBASE, EMBAL, and LEMBASE reloaded with enhancements
NEWS 14 SEP 24
                 CA/CAplus enhanced with pre-1907 records from Chemisches
NEWS 15 OCT 02
                 Zentralblatt
NEWS 16 OCT 19
                 BEILSTEIN updated with new compounds
NEWS 17 NOV 15 Derwent Indian patent publication number format enhanced
NEWS 18 NOV 19
                 WPIX enhanced with XML display format
NEWS 19 NOV 30 ICSD reloaded with enhancements
NEWS 20 DEC 04 LINPADOCDB now available on STN
NEWS 21 DEC 14 BEILSTEIN pricing structure to change
NEWS 22 DEC 17
                 USPATOLD added to additional database clusters
NEWS 23 DEC 17
                 IMSDRUGCONF removed from database clusters and STN
NEWS 24 DEC 17
                 DGENE now includes more than 10 million sequences
NEWS 25 DEC 17
                 TOXCENTER enhanced with 2008 MeSH vocabulary in
                 MEDLINE segment
         DEC 17
                 MEDLINE and LMEDLINE updated with 2008 MeSH vocabulary
NEWS 26
         DEC 17
NEWS 27
                 CA/CAplus enhanced with new custom IPC display formats
         DEC 17
                 STN Viewer enhanced with full-text patent content
NEWS 28
                 from USPATOLD
NEWS EXPRESS 19 SEPTEMBER 2007: CURRENT WINDOWS VERSION IS V8.2,
              CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
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AND CURRENT DISCOVER FILE IS DATED 19 SEPTEMBER 2007.

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Young, Shawquia, Page 1

NEWS HOURS

NEWS IPC8 For general information regarding STN implementation of IPC 8

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=> file reg COST IN U.S. DOLLARS

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STRUCTURE FILE UPDATES: 30 DEC 2007 HIGHEST RN 959750-30-2 DICTIONARY FILE UPDATES: 30 DEC 2007 HIGHEST RN 959750-30-2

New CAS Information Use Policies, enter HELP USAGETERMS for details.

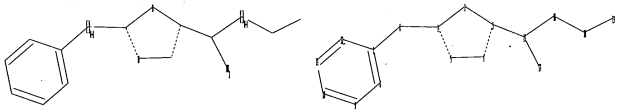
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http://www.cas.org/support/stngen/stndoc/properties.html

Uploading C:\Program Files\Stnexp\Queries\10578216.str



chain nodes :

6 15 16 17 18 19

ring nodes :

1 2 3 4 5 7 8 9 10 11 12

Young, Shawquia, Page 2

chain bonds :

1-6 3-15 6-7 15-16 15-17 16-18 18-19

ring bonds :

1-2 1-5 2-3 3-4 4-5 7-8 7-12 8-9 9-10 10-11 11-12

exact/norm bonds :

1-2 1-5 2-3 3-4 4-5 15-17

exact bonds :

1-6 3-15 6-7 15-16 16-18 18-19

normalized bonds :

7-8 7-12 8-9 9-10 10-11 11-12

isolated ring systems :

containing 1 : 7 :

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:CLASS 7:Atom 8:Atom 9:Atom 11:Atom 12:Atom 15:CLASS 16:GLASS 17:CLASS 18:CLASS 19:CLASS

L1 STRUCTURE UPLOADED

=> d l1

L1 HAS NO ANSWERS

L1 STR

Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 11:01:34 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 85 TO ITERATE

100.0% PROCESSED 85 ITERATIONS 0 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 1147 TO 2253

PROJECTED ANSWERS: 0 TO 0

L2 0 SEA SSS SAM L1

=> s l1 full

FULL SEARCH INITIATED 11:01:38 FILE 'REGISTRY'

FULL SCREEN SEARCH COMPLETED - 1463 TO ITERATE

100.0% PROCESSED 1463 ITERATIONS 6 ANSWERS

Young, Shawquia, Page 3

SEARCH TIME: 00.00.01

L3 6 SEA SSS FUL L1

=> file hcaplus
COST IN U.S. DOLLARS

SINCE FILE TOTAL SESSION 172.10 172.31

FULL ESTIMATED COST

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This file contains CAS Registry Numbers for easy and accurate substance identification.

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=> d ed abs ibib hitstr tot

ANSWER 1 OF 3 HCAPLUS COPYRIGHT 2007 ACS on STN Entered STN: 02 May 2006

Nonracemic imidazolemethanamines I (R = Me, Me2CH, Me2CHCH2, PhCH2) and

(R = Me2CH, Me2CHCH2, PhCH2, Rl = 2-pyridinyl, 2-HOC6H4) are prepared as tridentate ligands for copper-catalyzed Henry reactions of nitromethane with benzaldehyde or 4-nitrobenzaldehyde. Reductive amination of nonracemic imidazolemethanamines with aldehydes using either palladium-catalyzed hydrogenation or condensation and reduction of the

intermediates yields the title compds. The ligands prepared exhibit

strong
hydrogen bonding in d6-DMSO solution, resulting in hindered imidazole
tautomerism. In the presence of I (R = Me, Me2CH, Me2CHCH2, PhCH2) and

(R = Me2CH, Me2CHCH2, PhCH2; R1 = 2-pyridinyl, 2-HOC6H4) and copper (II) salts such as copper (II) 4-methoxybenzoate, benzaldehyde or 4-nitrobenzaldehyde undergo Henry reactions with nitromethane to give (S)-4-02NC6H4CH(OH2CNZO) or (S)-PhCH(OH)CHZNO2 (Or, in one case, (R)-4-02NC6H4CH(OH)CHZNO2) in 54-964 yields and in 8-324 ee. The Henry reactions of nitromethane and either benzaldehyde or 4-nitrobenzaldehyde in the presence of nonracemic imidazolemethanamines and copper (II) salts is optimized on temperature, stoichiometry, solvent, and the copper Ce. source.

Attempted Henry reaction of acetophenone and nitromethane in the presence of a copper (II) salt and a nonracemic bidentate imidazolemethanamine gare years of a copper (II) salt and a nonracemic bidentate imidazolemethanamine gives no product.

ACCESSION NUMBER: 2006:398391 HCAPLUS
DOCUMENT NUMBER: 145:83270

2006:398391 HCAPLUS
145:83270
Novel nitrogen ligands based on imidazole derivatives
and their application in asymmetric catalysis
Bures, Filip, Szotkowski, Tomas, Kulhanek, Jiri;
Pytela, Oldrich, Ludwig, Miroslav, Holcapek, Michal
Department of Organic Chemistry, University of
Pardubice, Pardubice, 53210, Czech Rep.
Tetrahedron: Asymmetry (2006), 17(6), 900-907
CODEN: TASYES; ISSN: 0957-4166 AUTHOR (S):

CORPORATE SOURCE:

SOURCE:

PUBLISHER: Blsevier B.V.

DOCUMENT TYPE: LANGUAGE:

Journal English CASREACT 145:83270 OTHER SOURCE(S):

852248-41-0 RL: CAT (Catalyst use); USES (Uses)

ANSWER 1 OF 3 HCAPLUS COPYRIGHT 2007 ACS on STN (Continued) ANSMER 1 OF 3 HCAPLUS COPYRIGHT 2007 ACS on STN (Continued)

(comparison of nonracemic imidazolemethanamines and a nonracemic imidazolylpyrrolidine as ligands in copper-catalyzed enanticselective Henry reactions of benzaldehydes and nitromethane)

852248-41-0 HCAPLUS

HI-Imidazole-4-methanamine, a-[(18)-1-methylpropy1]-2-pheny1-,

(aS)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

IT 852248-38-5 852248-40-9
RL: CAT (Catalyst use); RCT (Reactant); RACT (Reactant or reagent); USES

(preparation of nonracemic imidazolemethanamines by reductive amination of

ation of nonracemic imidazolemethanamines with aldehydes and their use as ligands for copper-catalyzed enantioselective Henry reactions) 852248-38-5 HCAPLUS H-Imidazole-4-methanamine, α-(1-methylethyl)-2-phenyl-, (αS)-(9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

852248-40-9 HCAPLUS 1H-Imidazole-4-methanamine, α -(2-methylpropyl)-2-phenyl-, (α S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

THERE ARE 14 CITED REFERENCES AVAILABLE FOR REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L4 ANSWER 2 OF 3 HCAPLUS COPYRIGHT 2007 ACS ON STN ED Entered STN: 28 Mar 2005 AB A route to the preparation of enantiopure ligands based on a 2-phenylindszole

nylimidazole ring is described. The stereogenic center is placed into the chain

ring is described. The stereogenic center is placed into the chain bonded to the fourth carbon of the imidazole ring. The synthesis starts from inexpensive and readily available N-protected \(\alpha\) and a subsequently, into \(\alpha\) becomes the source of chirality, which are converted into appropriate \(\alpha\) discount of the source and, subsequently, into \(\alpha\) becomes. These \(\alpha\) becomes a form ketones are good precursors for reactions with amidines to provide the imidazole ring. The deprotection into the final products was carried out using hydrogen.

ACCESSION NUMBER: 2005:263751 HCADLUS

DOCUMENT NUMBER: 143:7647

ITILE: Chiral imidazole derivatives synthesis from enantiopure N-protected \(\alpha\) amino \(\alpha\) enantiopure N-protected \(\alpha\) enantiopure N-protected \(\alpha\) amino \(\alpha\) enantiopure N-protected \(\alph

PUBLISHER: Slevier B.V.
DOCUMENT TYPE: Journal
LANGUAGE: English
OTHER SOURCE(S): CASREACT 143:7647

IT 852248-38-5P 852248-40-9P.852248-41-0P
RL: SPN (Synthetic preparation), PREP (Preparation)
(preparation of chiral imidazole derive, from enantiopure N-protected
a-amino acids)

RN 852248-38-5 NCAPLUS
CN 1H-Imidazole-4-methanamine, a-(1-methylethyl)-2-phenyl-, (aS)(9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

852248-40-9 HCAPLUS
1H-Imidazole-4-methanamine, α-(2-methylpropyl)-2-phenyl-,
(αs)- (9C1) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

852248-41-0 HCAPLUS
1H-Imidazole-4-methanamine, α -[(1S)-1-methylpropyl]-2-phenyl-,

ANSWER 2 OF 3 HCAPLUS COPYRIGHT 2007 ACS ON STN (GS) - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

REFERENCE COUNT:

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L4 ANSWER 3 OF 3 HCAPLUS COPYRIGHT 2007 ACS on STN (Continued)
Suoming; Zhao, He; Chenard, Bertrand L.; Gao, Yang;
Han, Bingsong; He, Xiao Shu
Neurogen Corporation, USA
POT Int. Appl., 356 pp.
CODEN; PIXXD2

DOCUMENT TYPE: - Batent
LANGUAGE: Brill ACC. NUM. COUNT: 1 FAMILY ACC. NUM. COUNT:

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US	2007	2080	48		A1		2007	0906		US	2007-	6808	65		- 2	20070	30.
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										US	2002-	3921	45P		Р :	20020	62
										US	2003 -	4059	8 9		A3 2	20030	32
											2003-						

OTHER SOURCE(S): MARPAT 139:307760
IT 610286-31-2P
RL: RCT (Reactant), SPN (Synthetic preparation), PREP (Preparation), RACT
(Reactant or reagent)
(preparation of new aryl imidazoles and related compds. as C5a
receptor

ptor modulators) 610286-31-2 HCAPLUS 1H-Enidacole-5-methanamine, a,1-dibutyl-2-phenyl-, (aR)- (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 3 OF 3 HCAPLUS COPYRIGHT 2007 ACS ON STN Entered STN: 10 Oct 2003

The title imidazoles, pyrazoles, pyridizines [I, the ring system in the formula I = 5-membered heteroaryl ring system (in which x=0, A=C, N, O, S, and E and G=C, N, provided that the 5-membered heteroaryl ring system does not contain more than 3 heteroatoms or more than 1 O or S atom) or 6-membered heteroaryl ring system (in which x=1, A, B, E, and AΒ

= C, N, and provided that the 6-membered heteroaryl ring system does not contain more than 3 N atoms); R, Rl = H, OH, halo, etc., when E = N, the R2 = alkyl, alkenyl, CH2Ph, etc., when E = C, then R2 = H, halo, OH,

, R3 = H, alkyl, alkenyl, etc.; R4 = alkyl, alkenyl, cycloalkyl, etc.; R5, R6 = H, alkyl; z = 1-3; Ar1 = (un)substituted aryl, heteroaryl, Ph fused to 5-7 membered (un)saturated ring that has 0-2 ring atoms chosen from

and S, Ar2 - cycloalkyl, cycloalkylalkyl, aryl having 1 ring or 2 fused

pendant rings, etc.; y = 1-6) which are ligands of C5a receptors, were prepared and formulated. E.g., a multi-step synthesis of II (starting

from

Me benzimidate hydrochloride and 1-butylamine), was given. Preferred compds. I bind to C5a receptors with high affinity (biol. data given) and exhibit neutral antagonist or inverse agonist activity at C5a receptors. This invention also relates to pharmaceutical compns. comprising such compds. It further relates to the use of such compds. in treating a variety of inflammatory and immune system disorders.

ACCESSION NUMBER: 2003:796668 HCAPLUS
DOCUMENT NUMBER: 139:307760
Preparation of new aryl imidazoles and related compounds as C5a receptor modulators
INVENTOR(S): Luke, George P.; Maynard, George, Mitchell, Scott; Thurkauf, Andrew, Xie, Linghong, Zhang, Luyan, Zhang,

ANSWER 3 OF 3 HCAPLUS COPYRIGHT 2007 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

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FOLD ESTIMATED COST	10.41	150.72
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION

-2.34 -2.34

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